Package ‘bayesdfa’

September 28, 2021

Type Package

Title Bayesian Dynamic Factor Analysis (DFA) with 'Stan'

Version 1.2.0

Description Implements Bayesian dynamic factor analysis with 'Stan'. Dynamic factor analysis is a dimension reduction tool for multivariate time series. 'bayesdfa' extends conventional dynamic factor models in several ways. First, extreme events may be estimated in the latent trend by modeling process error with a student-t distribution. Second, alternative constraints (including proportions are allowed). Third, the estimated dynamic factors can be analyzed with hidden Markov models to evaluate support for latent regimes.

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Encoding UTF-8

Depends R (>= 3.5.0)

Imports dplyr, ggplot2, loo (>= 2.0.0), methods, mgcv (>= 1.8.13), Rcpp (>= 0.12.0), RcppParallel (>= 5.0.1), reshape2, rlang, rstan (>= 2.18.1), rstantools (>= 2.1.1), viridisLite

LinkingTo BH (>= 1.66.0), Rcpp (>= 0.12.0), RcppEigen (>= 0.3.3.3.0), RcppParallel (>= 5.0.1), rstan (>= 2.18.1), StanHeaders (>= 2.18.1)

Suggests testthat, parallel, knitr, rmarkdown

URL https://fate-ewi.github.io/bayesdfa/

BugReports https://github.com/fate-ewi/bayesdfa/issues

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VignetteBuilder knitr

SystemRequirements GNU make

Biarch true

NeedsCompilation yes
R topics documented:

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The 'bayesdfa' package.

Description

A DESCRIPTION OF THE PACKAGE

References


dfa_cv

Apply cross validation to DFA model

Description

Apply cross validation to DFA model

Usage

dfa_cv(
    stanfit,
    cv_method = c("loocv", "lfocv"),
    fold_ids = NULL,
    n_folds = 10,
    iter = 2000,
    chains = 4,
    thin = 1,
    ...
)

Arguments

- **stanfit**
  A stanfit object, to preserve the model structure from a call to fit_dfa()
- **cv_method**
  The method used for cross validation. The options are 'loocv', where time is ignored and each data point is assigned randomly to a fold. The method 'lfocv' is leave time out cross validation, and time slices are iteratively held out out. Finally the method 'lfocv' implements leave future out cross validation to do one-step ahead predictions.
- **fold_ids**
  A vector whose length is the same as the number of total data points. Elements are the fold id of each data point. If not all data points are used (e.g. the lfocv or ltocv approach might only use 10 time steps) the value can be something other than a number, e.g. NA
### Examples

```r
# Not run:
set.seed(42)
s <- sim_dfa(num_trends = 1, num_years = 20, num_ts = 3)
obs <- c(s$y_sim[1, ], s$y_sim[2, ], s$y_sim[3, ])
long <- data.frame("obs" = obs, "ts" = sort(rep(1:3, 20)),
"time" = rep(1:20, 3))
m <- fit_dfa(y = long, iter = 50, chains = 1, data_shape = "long",
sample = FALSE)
  # random folds
fit_cv <- dfa_cv(m, cv_method = "loocv", n_folds = 5, iter = 50, chains = 1)

  # folds can also be passed in
fold_ids <- sample(1:5, size = nrow(long), replace = TRUE)
m <- fit_dfa(y = long, iter = 50, chains = 1, data_shape = "long",
sample = FALSE)
fit_cv <- dfa_cv(m, cv_method = "loocv", n_folds = 5, iter = 50, chains = 1,
fold_ids = fold_ids)

  # do an example of leave-time-out cross validation where years are dropped
fold_ids <- long$time
m <- fit_dfa(y = long, iter = 50, chains = 1, data_shape = "long",
sample = FALSE)
fit_cv <- dfa_cv(m, cv_method = "loocv", iter = 100, chains = 1,
fold_ids = fold_ids)

  # example with covariates and long format data
obs_covar <- expand.grid("time" = 1:20, "timeseries" = 1:3,
"covariate" = 1:2)
obs_covar$value <- rnorm(nrow(obs_covar), 0, 0.1)
obs <- c(s$y_sim[1, ], s$y_sim[2, ], s$y_sim[3, ])
m <- fit_dfa(y = long, iter = 50, chains = 1, obs_covar = obs_covar,
data_shape = "long", sample = FALSE)
fit_cv <- dfa_cv(m, cv_method = "loocv", n_folds = 5, iter = 50, chains = 1)
```

## End(Not run)
**dfa_loadings**

Description

Get the loadings from a DFA as a data frame

Usage

dfa_loadings(rotated_modelfit, names = NULL, summary = TRUE, conf_level = 0.95)

Arguments

- **rotated_modelfit**: Output from `fit_dfa`.
- **names**: Optional vector of names for time series labels. Should be same length as the number of time series.
- **summary**: Logical indicating whether to include summary statistics.
- **conf_level**: Probability level for CI.

Value

A data frame with the following columns: ID is an identifier for each time series, time is the time step, y is the observed values standardized to mean 0 and unit variance, estimate is the mean fitted value, lower is the lower CI, and upper is the upper CI.

See Also

predicted plot_fitted fit_dfa

Examples

```r
y <- sim_dfa(num_trends = 2, num_years = 20, num_ts = 4)
m <- fit_dfa(y = y$y_sim, num_trends = 2, iter = 50, chains = 1)
fitted <- dfa_fitted(m)
```
dfatrends

Arguments

rotated_modelfit
Output from `rotate_trends`.

names
An optional vector of names for plotting the loadings.

summary
Logical. Should the full posterior densities be returned? Defaults to TRUE.

cnf_level
Confidence level for credible intervals. Defaults to 0.95.

Value

A data frame with the following columns: name is an identifier for each loading, trend is the trend for the loading, median is the posterior median loading, lower is the lower CI, upper is the upper CI, and prob_diff0 is the probability the loading is different than 0. When summary = FALSE, there is no lower or upper columns and instead there are columns chain and draw.

See Also

plot_loadings fit_dfa rotate_trends

Examples

```r
set.seed(42)
s <- sim_dfa(num_trends = 2, num_ts = 4, num_years = 10)
# only 1 chain and 180 iterations used so example runs quickly:
m <- fit_dfa(y = s$y_sim, num_trends = 2, iter = 50, chains = 1)
r <- rotate_trends(m)
loadings <- dfa_loadings(r, summary = TRUE)
loadings <- dfa_loadings(r, summary = FALSE)
```

dfa_trends

Get the trends from a DFA as a data frame

Description

Get the trends from a DFA as a data frame

Usage

dfa_trends(rotated_modelfit, years = NULL)

Arguments

rotated_modelfit
Output from `rotate_trends`.

years
Optional numeric vector of years.
find_dfa_trends

Value
A data frame with the following columns: time is the time step, trend_number is an identifier for each trend, estimate is the trend mean, lower is the lower CI, and upper is the upper CI.

See Also
plot_trends fit_dfa rotate_trends

Examples
set.seed(1)
s <- sim_dfa(num_trends = 1)
m <- fit_dfa(y = s$y_sim, num_trends = 1, iter = 50, chains = 1)
r <- rotate_trends(m)
trends <- dfa_trends(r)

find_dfa_trends
Find the best number of trends according to LOOIC

Description
Fit a DFA with different number of trends and return the leave one out (LOO) value as calculated by the loo package.

Usage
find_dfa_trends(
  y = y,
  kmin = 1,
  kmax = 5,
  iter = 2000,
  thin = 1,
  compare_normal = FALSE,
  convergence_threshold = 1.05,
  variance = c("equal", "unequal"),
  ...
)

Arguments
y A matrix of data to fit. Columns represent time element.
kmin Minimum number of trends, defaults to 1.
kmax Maximum number of trends, defaults to 5.
iter Iterations when sampling from each Stan model, defaults to 2000.
thin Thinning rate when sampling from each Stan model, defaults to 1.
compare_normal If TRUE, does model selection comparison of Normal vs. Student-t errors
convergence_threshold
    The maximum allowed value of Rhat to determine convergence of parameters

variance
    Vector of variance arguments for searching over large groups of models. Can be either or both of ("equal","unequal")

... Other arguments to pass to `fit_dfa()`

Examples

```r
set.seed(42)
s <- sim_dfa(num_trends = 2, num_years = 20, num_ts = 3)
# only 1 chain and 180 iterations used so example runs quickly:
m <- find_dfa_trends(
y = s$y_sim, iter = 50,
kmin = 1, kmax = 2, chains = 1, compare_normal = FALSE,
    variance = "equal", convergence_threshold = 1.1,
    control = list(adapt_delta = 0.95, max_treedepth = 20)
)
m$summary
m$best_model
```

---

**find_inverted_chains**  
*Find which chains to invert*

**Description**
Find which chains to invert by checking the sum of the squared deviations between the first chain and each other chain.

**Usage**
```
find_inverted_chains(model, trend = 1, plot = FALSE)
```

**Arguments**
- **model**: A Stan model, `rstanfit` object
- **trend**: Which trend to check
- **plot**: Logical: should a plot of the trend for each chain be made? Defaults to `FALSE`

**See Also**
invert_chains
**find_regimes**

Fit multiple models with differing numbers of regimes to trend data

**Usage**

```r
find_regimes(
  y,
  sds = NULL,
  min_regimes = 1,
  max_regimes = 3,
  iter = 2000,
  thin = 1,
  chains = 1,
  ...
)
```

**Arguments**

- `y` Data, time series or trend from fitted DFA model.
- `sds` Optional time series of standard deviations of estimates. If passed in, residual variance not estimated.
- `min_regimes` Smallest of regimes to evaluate, defaults to 1.
- `max_regimes` Biggest of regimes to evaluate, defaults to 3.
- `iter` MCMC iterations, defaults to 2000.
- `thin` MCMC thinning rate, defaults to 1.
- `chains` MCMC chains; defaults to 1 (note that running multiple chains may result in a "label switching" problem where the regimes are identified with different IDs across chains).
- `...` Other parameters to pass to `rstan::sampling()`.

**Examples**

```r
data(Nile)
find_regimes(log(Nile), iter = 50, chains = 1, max_regimes = 2)
```
find_swans

Find outlying "black swan" jumps in trends

Description

Find outlying "black swan" jumps in trends

Usage

find_swans(rotated_modelfit, threshold = 0.01, plot = FALSE)

Arguments

- `rotated_modelfit`: Output from `rotate_trends()`.  
- `threshold`: A probability threshold below which to flag trend events as extreme  
- `plot`: Logical: should a plot be made?

Value

Prints a ggplot2 plot if `plot = TRUE`; returns a data frame indicating the probability that any given point in time represents a "black swan" event invisibly.

References


Examples

```r
set.seed(1)
s <- sim_dfa(num_trends = 1, num_ts = 3, num_years = 30)
s$s_y_sim[1, 15] <- s$s_y_sim[1, 15] - 6
plot(s$s_y_sim[1, ], type = "o")
abline(v = 15, col = "red")
# only 1 chain and 250 iterations used so example runs quickly:
m <- fit_dfa(y = s$s_y_sim, num_trends = 1, iter = 50, chains = 1, nu_fixed = 2)
r <- rotate_trends(m)
p <- plot_trends(r)  
#+ geom_vline(xintercept = 15, colour = "red")
print(p)
# a 1 in 1000 probability if was from a normal distribution:
find_swans(r, plot = TRUE, threshold = 0.001)
```
fit_dfa  

Fit a Bayesian DFA

Description

Fit a Bayesian DFA

Usage

```r
fit_dfa(
  y = y,
  num_trends = 1,
  varIndx = NULL,
  scale = c("zscore", "center", "none"),
  iter = 2000,
  chains = 4,
  thin = 1,
  control = list(adapt_delta = 0.99, max_treedepth = 20),
  nu_fixed = 101,
  est_correlation = FALSE,
  estimate_nu = FALSE,
  estimate_trend_ar = FALSE,
  estimate_trend_ma = FALSE,
  estimate_process_sigma = FALSE,
  equal_process_sigma = TRUE,
  estimation = c("sampling", "optimizing", "vb", "none"),
  data_shape = c("wide", "long"),
  obs_covar = NULL,
  pro_covar = NULL,
  z_bound = NULL,
  z_model = c("dfa", "proportion"),
  trend_model = c("rw", "bs", "ps", "gp"),
  n_knots = NULL,
  knot_locs = NULL,
  par_list = NULL,
  family = "gaussian",
  verbose = FALSE,
  gp_theta_prior = c(3, 1),
  expansion_prior = FALSE,
  ...
)
```

Arguments

- **y**: A matrix of data to fit. See `data_shape` option to specify whether this is long or wide format data. Wide format data (default) is a matrix with time across columns and unique time series across rows, and can only contain 1 observation.
fit_dfa

per time series - time combination. In contrast, long format data is a data frame that includes observations ("obs"), time ("time") and time series ("ts") identifiers – the benefit of long format is that multiple observations per time series can be included. Correlation matrix currently not estimated if data shape is long.

num_trends Number of trends to fit.

varIndx Indices indicating which timeseries should have shared variances.

scale Character string, used to standardized data. Can be "zscore" to center and standardize data, "center" to just standardize data, or "none". Defaults to "zscore"

iter Number of iterations in Stan sampling, defaults to 2000. Used for both rstan::sampling() and rstan::vb()

chains Number of chains in Stan sampling, defaults to 4.

thin Thinning rate in Stan sampling, defaults to 1.

control A list of options to pass to Stan sampling. Defaults to list(adapt_delta = 0.99, max_treedepth = 20).

nu_fixed Student t degrees of freedom parameter. If specified as greater than 100, a normal random walk is used instead of a random walk with a t-distribution. Defaults to 101.

est_correlation Boolean, whether to estimate correlation of observation error matrix $R$. Defaults to FALSE. Currently can’t be estimated if data are in long format.

estimate_nu Logical. Estimate the student t degrees of freedom parameter? Defaults to FALSE,

estimate_trend_ar Logical. Estimate AR(1) parameters on DFA trends? Defaults to ‘FALSE”, in which case AR(1) parameters are set to 1

estimate_trend_ma Logical. Estimate MA(1) parameters on DFA trends? Defaults to ‘FALSE“, in which case MA(1) parameters are set to 0.

estimate_process_sigma Logical. Defaults FALSE, whether or not to estimate process error sigma. If not estimated, sigma is fixed at 1, like conventional DFAs.

equal_process_sigma Logical. If process sigma is estimated, whether or not to estimate a single shared value across trends (default) or estimate equal values for each trend

estimation Character string. Should the model be sampled using rstan::sampling() ("sampling",default), rstan::optimizing() ("optimizing"), variational inference rstan::vb() ("vb"), or no estimation done ("none"). No estimation may be useful for debugging and simulation.

data_shape If wide (the current default) then the input data should have rows representing the various timeseries and columns representing the values through time. This matches the MARSS input data format. If long then the long format data is a data frame that includes observations ("obs"), time ("time") and time series ("ts") identifiers – the benefit of long format is that multiple observations per time series can be included
Optional dataframe of data with 4 named columns ("time","timeseries","covariate","value"), representing: (1) time, (2) the time series affected, (3) the covariate number for models with more than one covariate affecting each trend, and (4) the value of the covariate

Optional dataframe of data with 4 named columns ("time","trend","covariate","value"), representing: (1) time, (2) the trend affected, (3) the covariate number for models with more than one covariate affecting each trend, and (4) the value of the covariate

Optional hard constraints for estimated factor loadings – really only applies to model with 1 trend. Passed in as a 2-element vector representing the lower and upper bound, e.g. (0, 100) to constrain positive

Optional argument allowing for elements of Z to be constrained to be proportions (each time series modeled as a mixture of trends). Arguments can be "dfa" (default) or "proportion"

Optional argument to change the model of the underlying latent trend. By default this is set to 'rw', where the trend is modeled as a random walk - as in conventional DFA. Alternative options are 'bs', where B-splines are used to model the trends, "ps" where P-splines are used to model the trends, or 'gp', where gaussian predictive processes are used. If models other than 'rw' are used, there are some key points. First, the MA and AR parameters on these models will be turned off. Second, for B-splines and P-splines, the process_sigma becomes an optional scalar on the spline coefficients, and is turned off by default. Third, the number of knots can be specified (more knots = more wiggliness, and n_knots < N). For models with > 2 trends, each trend has their own spline coefficients estimated though the knot locations are assumed shared. If knots aren't specified, the default is N/3. By default both the B-spline and P-spline models use 3rd degree functions for smoothing, and include an intercept term. The P-spline model uses a difference penalty of 2.

The number of knots for the B-spline, P-spline, or Gaussian predictive process models. Optional, defaults to round(N/3)

Locations of knots (optional), defaults to uniform spacing between 1 and N

A vector of parameter names of variables to be estimated by Stan. If NULL, this will default to c("x", "Z", "sigma", "log_lik", "psi","xstar") for most models – though if AR / MA, or Student-t models are used additional parameters will be monitored. If you want to use diagnostic tools in rstan, including moment_matching, you will need to pass in a larger list. Setting this argument to "all" will monitor all parameters, enabling the use of diagnostic functions – but making the models a lot larger for storage. Finally, this argument may be a custom string of parameters to monitor, e.g. c("x","sigma")

String describing the observation model. Default is "gaussian", but included options are "gamma", "lognormal", negative binomial ("nbinom2"), "poisson", or "binomial". The binomial family is assumed to have logit link, gaussian family is assumed to be identity, and the rest are log-link.

Whether to print iterations and information from Stan, defaults to FALSE.

A 2-element vector controlling the prior on the Gaussian process parameter in cov_exp_quad. This prior is a half-Student t prior, with the first argument of
fit_dfa

gp_theta_prior being the degrees of freedom (nu), and the second element being
the standard deviation

expansion_prior

Defaults to FALSE, if TRUE uses the parameter expansion prior of Ghosh &
Dunson 2009

... Any other arguments to pass to \texttt{rstan::sampling}().

Details

Note that there is nothing restricting the loadings and trends from being inverted (i.e. multiplied by
\(-1\)) for a given chain. Therefore, if you fit multiple chains, the package will attempt to determine
which chains need to be inverted using the function \texttt{find_inverted_chains}().

See Also

plot_loadings plot_trends rotate_trends find_swans

Examples

```r
set.seed(42)
s <- sim_dfa(num_trends = 1, num_years = 20, num_ts = 3)
# only 1 chain and 250 iterations used so example runs quickly:
m <- fit_dfa(y = s$y_sim, iter = 50, chains = 1)
## Not run:
# example of observation error covariates
set.seed(42)
obs_covar <- expand.grid("time" = 1:20, "timeseries" = 1:3, "covariate" = 1)
obs_covar$value <- rnorm(nrow(obs_covar), 0, 0.1)
m <- fit_dfa(y = s$y_sim, iter = 50, chains = 1, obs_covar = obs_covar)

# example of process error covariates
pro_covar <- expand.grid("time" = 1:20, "trend" = 1:2, "covariate" = 1)
pro_covar$value <- rnorm(nrow(pro_covar), 0, 0.1)
m <- fit_dfa(y = s$y_sim, iter = 50, chains = 1, num_trends = 2, pro_covar = pro_covar)

# example of long format data
s <- sim_dfa(num_trends = 1, num_years = 20, num_ts = 3)
obs <- c(s$y_sim[1, ], s$y_sim[2, ], s$y_sim[3, ])
long <- data.frame("obs" = obs, "ts" = sort(rep(1:3, 20)), "time" = rep(1:20, 3))
m <- fit_dfa(y = long, data_shape = "long", iter = 50, chains = 1)

# example of long format data with obs covariates
s <- sim_dfa(num_trends = 1, num_years = 20, num_ts = 3)
obs <- c(s$y_sim[1, ], s$y_sim[2, ], s$y_sim[3, ])
long <- data.frame("obs" = obs, "ts" = sort(rep(1:3, 20)), "time" = rep(1:20, 3))
obs_covar <- expand.grid("time" = 1:20, "timeseries" = 1:3, "covariate" = 1:2)
obs_covar$value <- rnorm(nrow(obs_covar), 0, 0.1)
m <- fit_dfa(y = long, data_shape = "long", iter = 50, chains = 1, obs_covar = obs_covar)

# example of model with Z constrained to be proportions and wide format data
s <- sim_dfa(num_trends = 1, num_years = 20, num_ts = 3)
m <- fit_dfa(y = s$y_sim, z_model = "proportion", iter = 50, chains = 1)
```
# example of model with Z constrained to be proportions and long format data
s <- sim_dfa(num_trends = 1, num_years = 20, num_ts = 3)
obs <- c(s$y_sim[1,], s$y_sim[2,], s$y_sim[3,])
long <- data.frame("obs" = obs, "ts" = sort(rep(1:3, 20)), "time" = rep(1:20, 3))
m <- fit_dfa(y = long, data_shape = "long", z_model = "proportion", iter = 50, chains = 1)

# example of B-spline model with wide format data
s <- sim_dfa(num_trends = 1, num_years = 20, num_ts = 3)
m <- fit_dfa(y = s$y_sim, iter = 50, chains = 1, trend_model = "bs", n_knots = 10)

# example of P-spline model with wide format data
s <- sim_dfa(num_trends = 1, num_years = 20, num_ts = 3)
m <- fit_dfa(y = s$y_sim, iter = 50, chains = 1, trend_model = "ps", n_knots = 10)

# example of Gaussian process model with wide format data
s <- sim_dfa(num_trends = 1, num_years = 20, num_ts = 3)
m <- fit_dfa(y = s$y_sim, iter = 50, chains = 1, trend_model = "gp", n_knots = 5)

## End(Not run)
invert_chains

chains MCMC chains, defaults to 1 (note that running multiple chains may result in a label switching problem where the regimes are identified with different IDs across chains).

... Other parameters to pass to rstan::sampling().

Examples

    data(Nile)
    fit_regimes(log(Nile), iter = 50, n_regimes = 1)

hmm_init Create initial values for the HMM model.

Description

Create initial values for the HMM model.

Usage

    hmm_init(K, x_t)

Arguments

K The number of regimes or clusters to fit. Called by rstan::sampling().

x_t A matrix of values. Called by rstan::sampling().

Value

list of initial values (mu, sigma)

invert_chains Invert chains

Description

Invert chains

Usage

    invert_chains(model, trends = 1, print = FALSE, ...)

Arguments

model A Stan model, rstanfit object

trends The number of trends in the DFA, defaults to 1

print Logical indicating whether the summary should be printed. Defaults to FALSE.

... Other arguments to pass to find_inverted_chains().
is_converged

**See Also**

find_inverted_chains

---

### is_converged

**Description**

Pass in rstanfit model object, and a threshold Rhat value for convergence. Returns boolean.

**Usage**

```r
is_converged(fitted_model, threshold = 1.05, parameters = c("sigma", "x", "Z"))
```

**Arguments**

- `fitted_model`: Samples extracted (with permuted = FALSE) from a Stan model. E.g. output from `invert_chains()`.
- `threshold`: Threshold for maximum Rhat.
- `parameters`: Vector of parameters to be included in convergence determination. Defaults = `c("sigma","x","Z")`. Other elements can be added including "pred", "log_lik", or "lp__"

---

### loo.bayesdfa

**Description**

Extract the LOOIC (leave-one-out information criterion) using `loo::loo()`. Note that we've implemented slightly different variants of loo, based on whether the DFA observation model includes correlation between time series or not (default is no correlation). Importantly, these different versions are not directly comparable to evaluate data support for including correlation or not in a DFA. If time series are not correlated, the point-wise log-likelihood for each observation is calculated and used in the loo calculations. However if time series are correlated, each time slice is assumed to be a joint observation of all variables, and the point-wise log-likelihood is calculated as the joint likelihood of all variables under the multivariate normal distribution.

**Usage**

```r
## S3 method for class 'bayesdfa'
loo(x, ...)
```

**Arguments**

- `x`: Output from `fit_dfa()`.
- `...`: Arguments for `loo::relative_eff()` and `loo::loo.array()`.
Examples

```
set.seed(1)
s <- sim_dfa(num_trends = 1, num_years = 20, num_ts = 3)
m <- fit_dfa(y = s$y_sim, iter = 50, chains = 1, num_trends = 1)
loo(m)
```

---

plot_fitted

Plot the fitted values from a DFA

Description

Plot the fitted values from a DFA

Usage

```
plot_fitted(
  modelfit,
  conf_level = 0.95,
  names = NULL,
  spaghetti = FALSE,
  time_labels = NULL
)
```

Arguments

- **modelfit**: Output from `fit_dfa`, a rstanfit object
- **conf_level**: Probability level for CI.
- **names**: Optional vector of names for plotting labels. Should be same length as the number of time series.
- **spaghetti**: Defaults to FALSE, but if TRUE puts all raw time series (grey) and fitted values on a single plot.
- **time_labels**: Optional vector of time labels for plotting, same length as number of time steps

See Also

plot_loadings fit_dfa rotate_trends dfa_fitted

Examples

```
y <- sim_dfa(num_trends = 2, num_years = 20, num_ts = 4)
m <- fit_dfa(y = y$y_sim, num_trends = 2, iter = 50, chains = 1)
p <- plot_fitted(m)
print(p)
```
p <- plot_fitted(m, spaghetti = TRUE)
print(p)

plot_loadings

Plot the loadings from a DFA

Description

Plot the loadings from a DFA

Usage

plot_loadings(
  rotated_modelfit,
  names = NULL,
  facet = TRUE,
  violin = TRUE,
  conf_level = 0.95,
  threshold = NULL
)

Arguments

rotated_modelfit
  Output from `rotate_trends()`.

names
  An optional vector of names for plotting the loadings.

facet
  Logical. Should there be a separate facet for each trend? Defaults to TRUE.

violin
  Logical. Should the full posterior densities be shown as a violin plot? Defaults to TRUE.

conf_level
  Confidence level for credible intervals. Defaults to 0.95.

threshold
  Numeric (0-1). Optional for plots, if included, only plot loadings who have Pr(<0) or Pr(>0) > threshold. For example threshold = 0.8 would only display estimates where 80% of posterior density was above/below zero. Defaults to NULL (not used).

See Also

plot_trends fit_dfa rotate_trends
Examples

```r
set.seed(42)
s <- sim_dfa(num_trends = 2, num_ts = 4, num_years = 10)
# only 1 chain and 180 iterations used so example runs quickly:
m <- fit_dfa(y = s$y_sim, num_trends = 2, iter = 50, chains = 1)
r <- rotate_trends(m)
plot_loadings(r, violin = FALSE, facet = TRUE)
plot_loadings(r, violin = FALSE, facet = FALSE)
plot_loadings(r, violin = TRUE, facet = FALSE)
plot_loadings(r, violin = TRUE, facet = TRUE)
```

---

**plot_regime_model**  
*Plot the state probabilities from* find_regimes(*)*

**Description**

Plot the state probabilities from find_regimes()

**Usage**

```r
plot_regime_model(
  model,
  probs = c(0.05, 0.95),
  type = c("probability", "means"),
  regime_prob_threshold = 0.9,
  plot_prob_indices = NULL,
  flip_regimes = FALSE
)
```

**Arguments**

- `model`  
  A model returned by find_regimes().
- `probs`  
  A numeric vector of quantiles to plot the credible intervals at.  
  Defaults to `c(0.05, 0.95)`.  
- `type`  
  Whether to plot the probabilities (default) or means.
- `regime_prob_threshold`  
  The probability density that must be above 0.5.  
  Defaults to 0.9 before we classify a regime (only affects "means" plot).
- `plot_prob_indices`  
  Optional indices of probability plots to plot.  
  Defaults to showing all.
- `flip_regimes`  
  Optional whether to flip regimes in plots, defaults to FALSE

**Details**

Note that the original timeseries data (dots) are shown scaled between 0 and 1.
Examples

```r
data(Nile)
m <- fit_regimes(log(Nile), n_regimes = 2, chains = 1, iter = 50)
plot_regime_model(m)
plot_regime_model(m, plot_prob_indices = c(2))
plot_regime_model(m, type = "means")
```

---

**plot_trends**

*Plot the trends from a DFA*

**Description**

Plot the trends from a DFA

**Usage**

```r
plot_trends(
  rotated_modelfit,
  years = NULL,
  highlight_outliers = FALSE,
  threshold = 0.01
)
```

**Arguments**

- `rotated_modelfit`  
  Output from `rotate_trends`
- `years`  
  Optional numeric vector of years for the plot
- `highlight_outliers`  
  Logical. Should trend events that exceed the probability of occurring with a normal distribution as defined by `threshold` be highlighted? Defaults to FALSE
- `threshold`  
  A probability threshold below which to flag trend events as extreme. Defaults to 0.01

**See Also**

`dfa_trends` `plot_loadings` `fit_dfa` `rotate_trends`

**Examples**

```r
set.seed(1)
s <- sim_dfa(num_trends = 1)
m <- fit_dfa(y = s$y_sim, num_trends = 1, iter = 50, chains = 1)
r <- rotate_trends(m)
p <- plot_trends(r)
print(p)
```
predicted

Calculate predicted value from DFA object

Description

Pass in `rstanfit` model object. Returns array of predictions, dimensioned number of MCMC draws x number of MCMC chains x time series length x number of time series

Usage

```
predicted(fitted_model)
```

Arguments

- `fitted_model`: Samples extracted (with `permuted` = FALSE) from a Stan model. E.g. output from `invert_chains()`.

Examples

```
## Not run:
set.seed(42)
s <- sim_dfa(num_trends = 1, num_years = 20, num_ts = 3)
# only 1 chain and 1000 iterations used so example runs quickly:
m <- fit_dfa(y = s$y_sim, iter = 2000, chains = 3, num_trends = 1)
pred <- predicted(m)
## End(Not run)
```

rotate_trends

Rotate the trends from a DFA

Description

Rotate the trends from a DFA

Usage

```
rotate_trends(fitted_model, conf_level = 0.95, invert = FALSE)
```

Arguments

- `fitted_model`: Output from `fit_dfa()`.
- `conf_level`: Probability level for CI.
- `invert`: Whether to invert the trends and loadings for plotting purposes
Examples

```r
set.seed(42)
s <- sim_dfa(num_trends = 1, num_years = 20, num_ts = 3)
# only 1 chain and 800 iterations used so example runs quickly:
m <- fit_dfa(y = s$y_sim, iter = 50, chains = 1)
r <- rotate_trends(m)
plot_trends(r)
```

---

**Description**

Simulate from a DFA

**Usage**

```r
sim_dfa(
  num_trends = 1,
  num_years = 20,
  num_ts = 4,
  loadings_matrix = matrix(nrow = num_ts, ncol = num_trends, rnorm(num_ts * num_trends, 0, 1)),
  sigma = rlnorm(1, meanlog = log(0.2), 0.1),
  varIndx = rep(1, num_ts),
  extreme_value = NULL,
  extreme_loc = NULL,
  nu_fixed = 100,
  user_supplied_deviations = NULL
)
```

**Arguments**

- **num_trends**  
The number of trends.
- **num_years**  
The number of years.
- **num_ts**  
The number of timeseries.
- **loadings_matrix**  
A loadings matrix. The number of rows should match the number of timeseries and the number of columns should match the number of trends. Note that this loadings matrix will be internally manipulated by setting some elements to 0 and constraining some elements to 1 so that the model can be fitted. See `fit_dfa()`.
See the outfit element Z in the returned list is to see the manipulated loadings matrix. If not specified, a random matrix $\sim N(0,1)$ is used.
- **sigma**  
A vector of standard deviations on the observation error. Should be of the same length as the number of trends. If not specified, random numbers are used `rlnorm(1, meanlog = log(0.2), 0.1)`.

varIndx  Indices of unique observation variances. Defaults to c(1,1,1,1). Unique observation error variances would be specified as c(1,2,3,4) in the case of 4 time series.

extreme_value  Value added to the random walk in the extreme time step. Defaults to not included.

extreme_loc  Location of single extreme event in the process. The same for all processes, and defaults to round(n_t/2) where n_t is the time series length

nu_fixed  Nu is the degrees of freedom parameter for the t-distribution, defaults to 100, which is effectively normal.

user_supplied_deviations  An optional matrix of deviations for the trend random walks. Columns are for trends and rows are for each time step.

Value  
A list with the following elements: y_sim is the simulated data, pred is the true underlying data without observation error added, x is the underlying trends, Z is the manipulated loadings matrix that is fed to the model.

Examples

```r
x <- sim_dfa(num_trends = 2)
names(x)
matplot(t(x$y_sim), type = "l")
matplot(t(x$x), type = "l")

set.seed(42)
x <- sim_dfa(extreme_value = -4, extreme_loc = 10)
matplot(t(x$x), type = "l")
abline(v = 10)
matplot(t(x$pred), type = "l")
abline(v = 10)

set.seed(42)
x <- sim_dfa()
matplot(t(x$x), type = "l")
abline(v = 10)
matplot(t(x$pred), type = "l")
abline(v = 10)
```

trend_cor  Estimate the correlation between a DFA trend and some other time-series

Description

Fully incorporates the uncertainty from the posterior of the DFA trend.
trend_cor

Usage

trend_cor(
  rotated_modelfit,
  y,
  trend = 1,
  time_window = seq_len(length(y)),
  trend_samples = 100,
  stan_iter = 300,
  stan_chains = 1,
  ...
)

Arguments

rotated_modelfit
  Output from rotate_trends().

y
  A numeric vector to correlate with the DFA trend. Must be the same length as the DFA trend.

trend
  A number corresponding to which trend to use, defaults to 1.

time_window
  Indices indicating a time window slice to use in the correlation. Defaults to using the entire time window. Can be used to walk through the timeseries and test the cross correlations.

trend_samples
  The number of samples from the trend posterior to use. A model will be run for each trend sample so this value shouldn’t be too large. Defaults to 100.

stan_iter
  The number of samples from the posterior with each Stan model run, defaults to 300.

stan_chains
  The number of chains for each Stan model run, defaults to 1.

...
  Other arguments to pass to sampling

Details

Uses a sigma ~ half_t(3,0,2) prior on the residual standard deviation and a uniform(-1,1) prior on the correlation coefficient. Fitted as a linear regression of y ~ x, where y represents the y argument to trend_cor() and x represents the DFA trend, and both y and x have been scaled by subtracting their means and dividing by their standard deviations. Samples are drawn from the posterior of the trend and repeatedly fed through the Stan regression to come up with a combined posterior of the correlation.

Value

A numeric vector of samples from the correlation coefficient posterior.

Examples

set.seed(1)
  s <- sim_dfa(num_trends = 1, num_years = 15)
  m <- fit_dfa(y = s$y_sim, num_trends = 1, iter = 50, chains = 1)
r <- rotate_trends(m)
n_years <- ncol(r$trends[, 1, ])  
fake_dat <- rnorm(n_years, 0, 1)
correlation <- trend_cor(r, fake_dat, trend_samples = 25)
hist(correlation)
correlation <- trend_cor(r,  
y = fake_dat, time_window = 5:15,  
trend_samples = 25)

hist(correlation)
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